Controlling immunity balances the brain in health and disease

Michal Schwartz

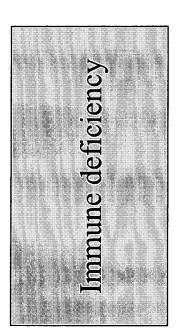
Department of Neurobiology

Homeostasis

Levels of immune activity



QuickTime™ and a TIFF (Uncompressed) decompressor are needed to see this picture.



are associated with a wide-spread Why neurodegenerative conditions loss of neurons:

- Poor spontaneous neurogenesis (limited formation of new neurons)
- Poor spontaneous regeneration (poor re-growth)
- · Diffuse damage due to high vulnerability to defense mechanism unless tightly controlled –'domino effect'

Common view of inflammation in neurodegenerative conditions

>In most neurodegenerative diseases there is a local inflammatory response. > This local inflammatory response (mediated by adaptive and/or innate immunity) has collectively received a bad reputation.

Our concept: The immune system plays a key role in Central Nervous System

maintenance

Renewal

Plasticity

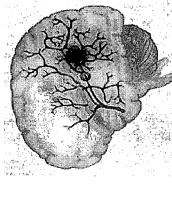
suppression over time denies the key players in We argue against bad or good innate/adaptive immunity or against good/bad cytokines in the be tightly controlled rather than suppressed context of the CNS; immune response should brain's maintenance and repair

Degenerative conditions and consequences of defense battle

Accumulating self compounds (e.g. gangliosides, B-amyloid, Prion, S-Ag, etc.)

And Single Singl

Growth Factors (Neurotrophins



otamate otamate

Legrond death

Consequences of defense

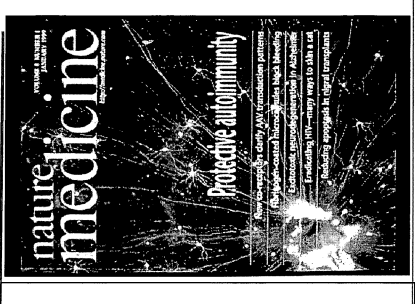
impodiance

Ioric

Malfunctioning microglia Inflammation-associated Toxicity (NO, TNF-a, COX-2)

Immune cells are needed for CNS maintenance and repair "Protective autoimmunity"

- Protective autoimmunity: A controlled T-cell response recognizing CNS antigens protects against internal enemies
- Autoimmune disease: An outcome of malfunctioning of autoimmunity
- Tolerance to self: Ability to tolerate response to self without developing an autoimmune disease
- Specificity provides the T cells with a way of homing and local reinforcement/activation.



Trends Immunol., 2002; Yoles et al., J. Neurosci. 2001; Kipnis et al., PNAS, 2001 IINS, 1999, 2003; Hauben et al. J. Neurosci., 2000, 2003; Schwartz and Kipnis, Rapalino et al., <u>Nat. Med</u>., 1998; Moalem et al., <u>Nat. Med</u>.1999; Schwartz et al., 2003; Kipnis et al., J. Neurosci., 2003; Mizrahi et al., J. Immunol., 2002.

myelin self-antigen prevents complete paralysis while Posttraumatic therapeutic vaccination with modified avoiding autoimmune disease

Ehud Hauben, Eugenia Agranov, Amalia Gothilf, Uri Nevo, Avi Cohen, Igor Smirnov, Lawrence Steinman, 3 and Michal Schwartz1

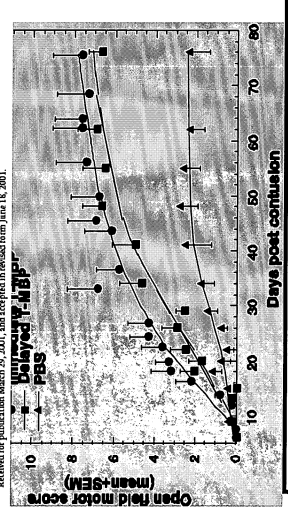
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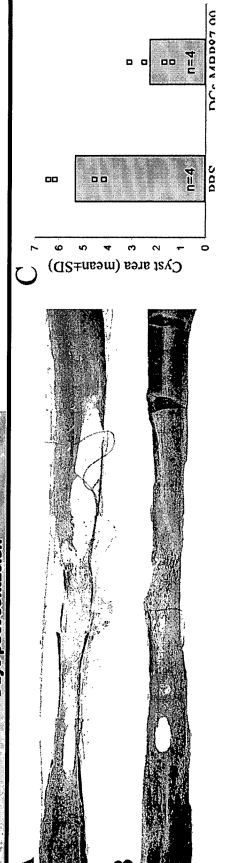
Proneuron Biorechnologies Ltd., Kiryat Weizmann, Ness Ziona, Israel

Department of Neurology, Stanford University School of Medicine, Stanford, California, USA

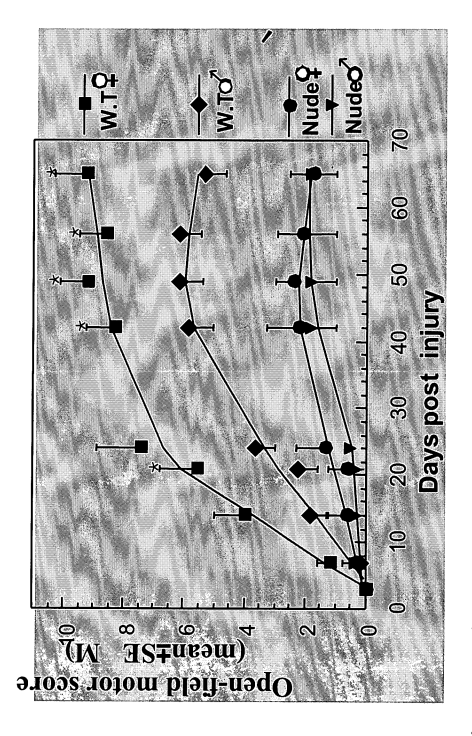
Address correspondence to: Michal Schwartz, Department of Neurobiology, The Weizmann Institute of Science, 76100 Rehovor, Israel. Phone: 972-8-93467; Fax 972-8-9344131; Email: michal.schwartz@weizmann.ac.il.

Received for publication March 29, 2001, and accepted in revised form June 18, 2001





Immune compromise animals have a weak ability to cope with spinal cord injury



Hauben et al., Eur. J. Neurosci., 2002

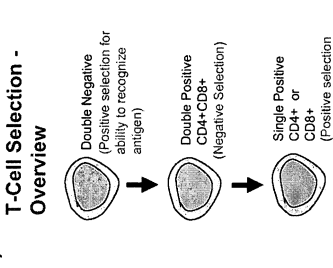


Recognizing Self

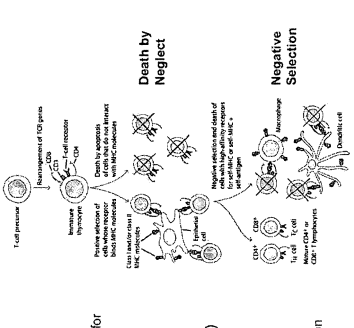
(nature mistake or

purposeful selection?) intruders)

(fighting against



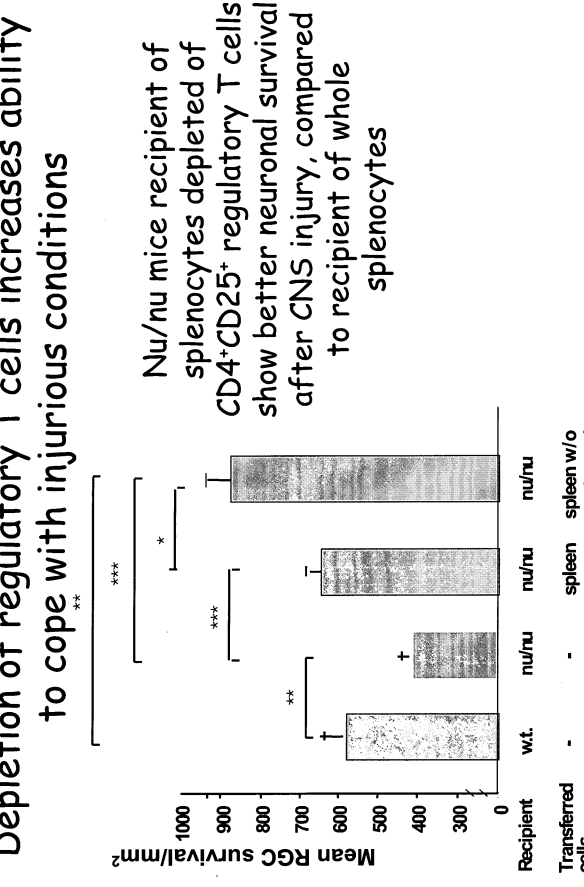
based on MHC reactivity)



What is the difference between beneficial and disease-causing autoimmune T cells? Affinity?

Number?

Persistence?

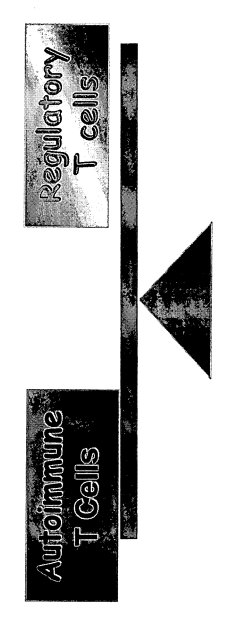


Kipnis et al., PNAS, 2002

CD4+CD25+

Signal (signal II) A.stress Signal is required to 'weaken' the What is needed to evoke autoimmunity? Effector T Cell CD4+CD25+ suppression Antigen Recognition Signal (Signal I) Co-Stimulatory Signal (signal II) Antigen Recognition Signal (Signal I) Local/Systemic Stress Signal from the Damaged Tissue (Signal III)

Balance Between Autoimmune T cells and Regulatory T cells



Autoimmune-dependent CNS Homeostasis

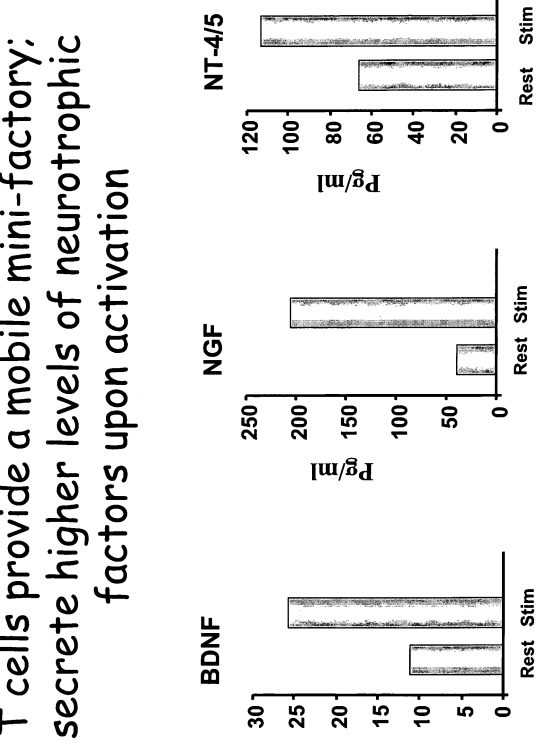
Malfunction

Too much Autoimmune disease

Too much
Neuronal loss
Limited neurogenesis

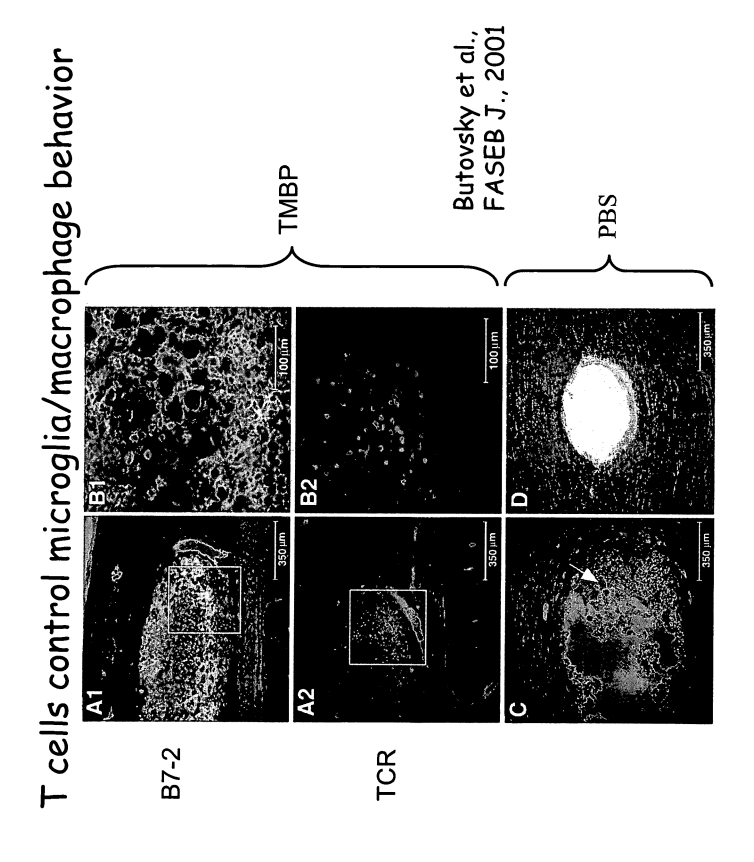
The underlying mechanism

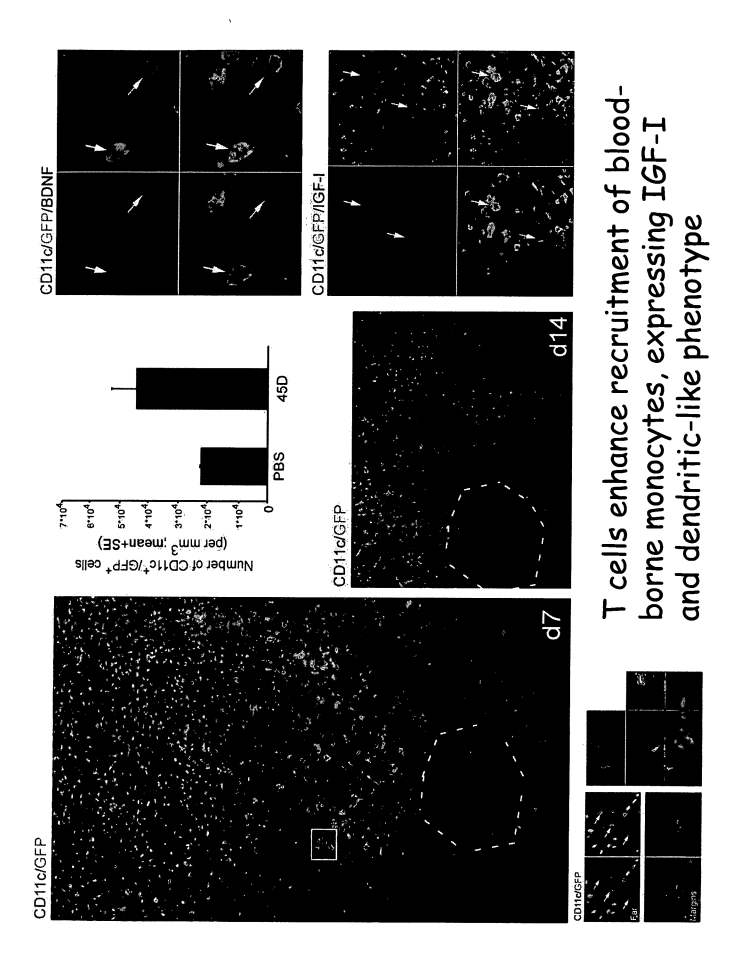
T cells provide a mobile mini-factory;



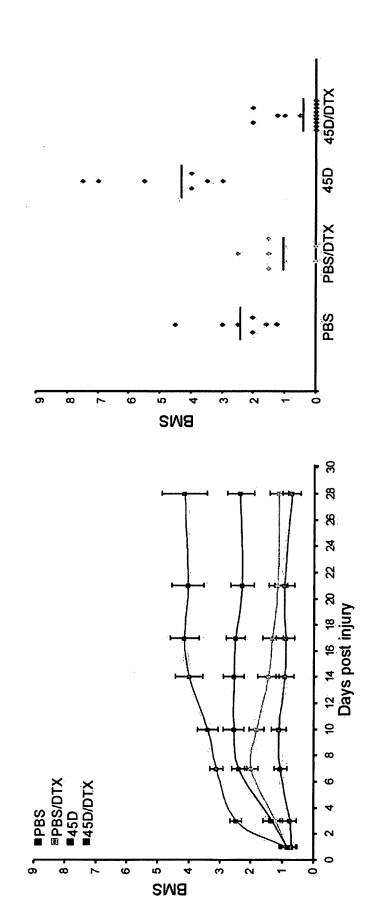
Pg/ml

Moalem et al., J. Autoimmun, 2000; Kipnis et al., PNAS, 2000



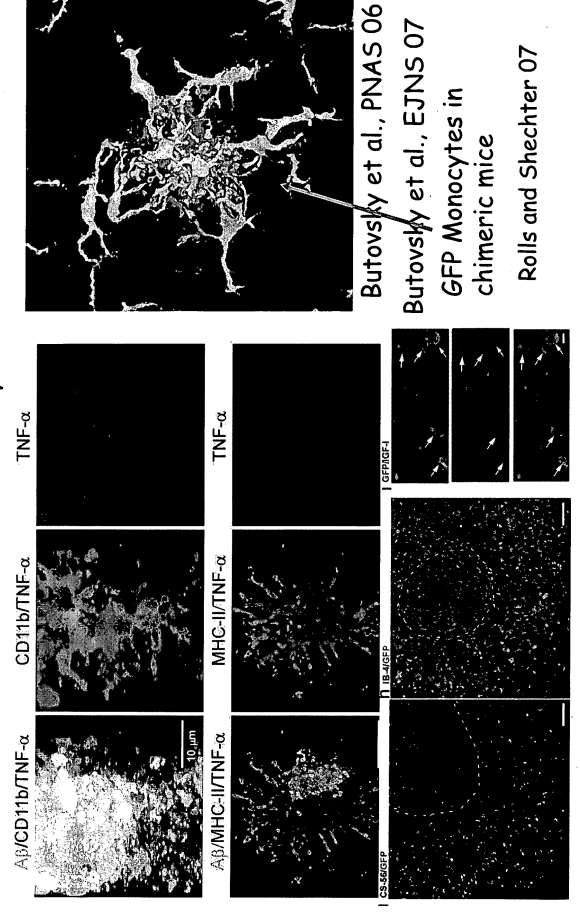


Selective depletion of bloodborne monocytes expressing CD11c completely impaired recovery XTQ oN



Depletion of CD11c+/GFP+ bone marrow-derived monocytes worsened recovery CX3CR1GFP No DTX

CNS specific T cells are needed for creating an immunological niche: microglial phenotype switch and recruitment of bloodborne monocytes



Microglia function as stand-by resident immune cells

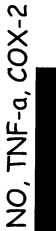
Bad

Neurodegeneration Innate activation



Immune functions -

killing and removal of microorganisms Secretion of:





poob

overwhelmed

Activation by adaptive immunity

Protection



Overwhelming activation:

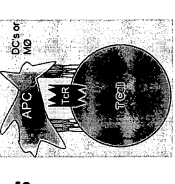
Immune and neural functions:

Delivery of neurotrophic

factors and cytokines

TNF-a

counteract the benefit Removal of growth inhibition



Buffering of toxicity mediators

(e.g. myelin phagocytosis)

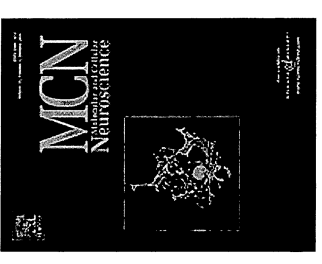
(e.g. Glutamate clearance)

Antigen presenting cells

The underlying mechanism

- cytokines and growth factors (Moalem et., Nat. Med. 1999; T cells serve as a mobile mini- factory producing locally J. Autoimmunity 2000; Kipnis et al., PNAS 2000)
- and BDNF, (b) act as antigen presenting cells (c) support T cells depending on their phenotypes 'shape' microglial activity and confer them with ability to (a) produce IGF-I neural tissue survival, (d) buffer glutamate, and (d) to support cell renewal
- (Butovsky et al., FASEB J., 2001; Mol. Cell. Neurosci., 2005, 2006; Shaked et al., <u>J.Neurochem.</u>, 2005; <u>J.Clin.</u> Inves., 2006; Ziv et al., Nat. Neurosci., 2006).

that the primary role of microglia is to maintain Working Hypothesis: Our in vitro findings that adult brain, their role in diseased conditions is neurogenesis/oligodendrogenesis in a healthy renewal from adult neural stem cells suggest T cell-activated microglia can support cell an extension of this primary role neuronal survival and



Butovsky, Ziv et al. Mol. Cell. Neurosci., 2005, 2006

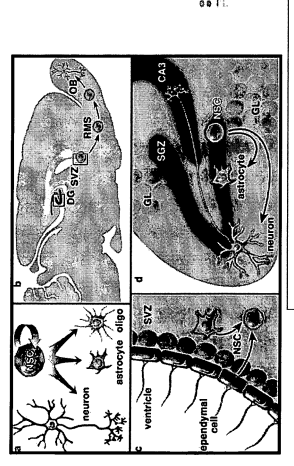
Cell renewal in the adult CNS

Nac Med 1958 North (1,1)1313-7

Neurogenesis in the adult human hippocampus.

Erksson PS. Perfilieva E. Bjork-Erksson I. Alborn AM. Nordborg C. Peterson D.A. Gage IH.

Neurogenesis at adulthood hippocampus and olfactory is restricted to

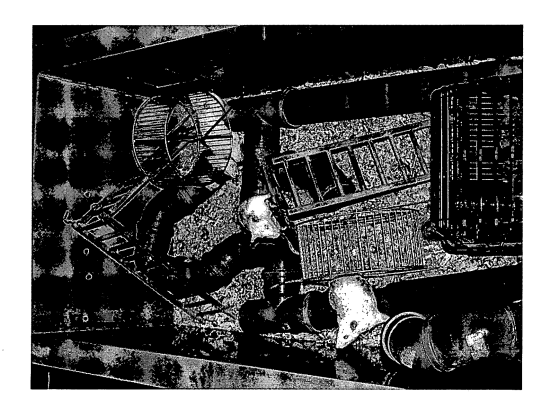


Stem cell proliferation and migration in Proliferation and differentiation of progenitor cells throughout the intact adult rat spinal cord. the intact adult spinal cord Honner KJ, Power AE, Kemperanan G, Kalin HG, Paliner ID, Winkler J. Thal LJ, Gage FH NEUROCENESIS MODELS Neurosci, 2000 Mar 15;20(6):2218-28.
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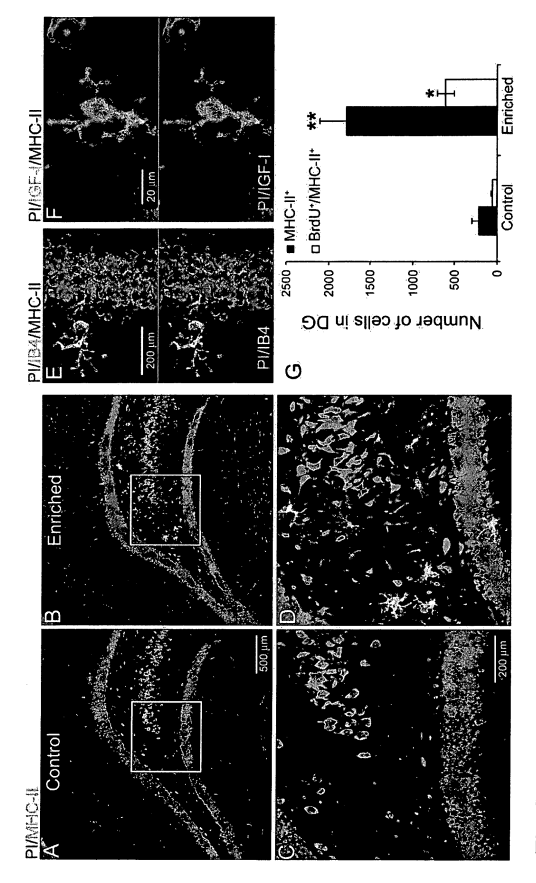
mammalian eye (Tropepe et al, Science 2000)

Retinal stem cells in the adult

Enriched Environment boosts formation of new neurons in the brain from a pool of adult stem cells



associated with microglia expressing a T-cell-activated Neurogenesis induced by enriched environment is phenotype



Ziv, Ron, Butovsky, et al., Nat. Neurosci., 2006

Are T cells contributing to the maintenance of the adult neurogenic niche?

Routes for circulating leukocytes to cross blood— CNS barriers

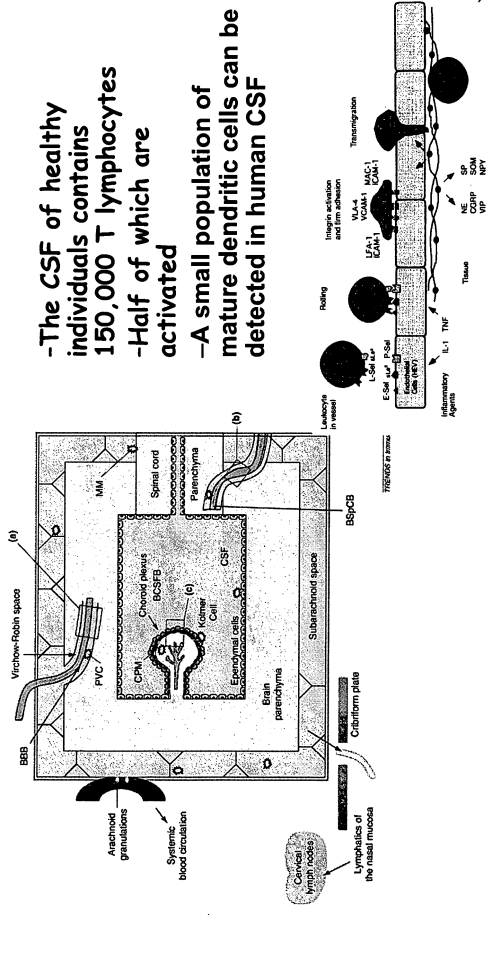
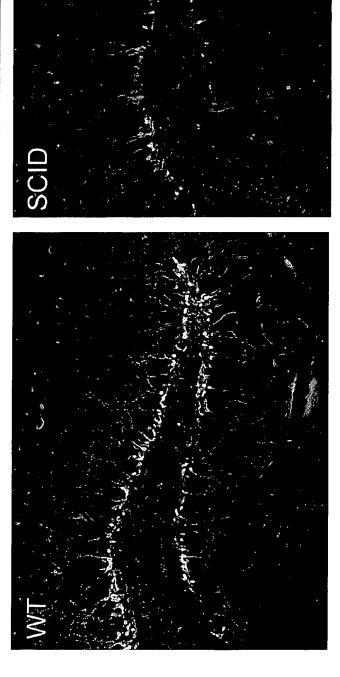
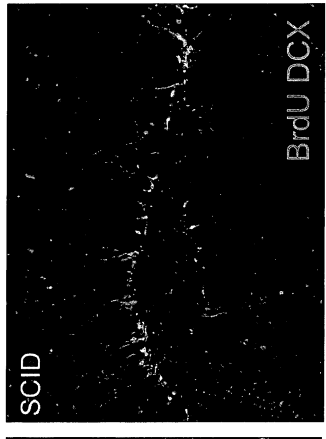


FIGURE 1 Leukocyte adhesion to endothelial cells and migration into tissues. Leukocyte adhesion and migration is a multistep process that initially involves (a) leukocyte rolling, (b) stimulation of integrins and firm adhesion, and (c) dispedesis into the tissues. Catecholamines such as norepinephrine (WE) and neuropeptides (calcitonin generaled peptide [CGRP], substance P[SP], sumatostatin [SOM], neuropeptide Y [NPT], vascactive intestant peptide [VIP] present in nerve terminals surrounding blood vessels or contained in the lissue parenchyma can modulate this process.

Adult Neurogenesis is impaired in immune deficient mice





Ziv, Ron, Butovsky et al., Nature Neurosci. 2006



SCID

₹

0 + CS781/6J

2,000

Cells per dentate gyrus

1,000

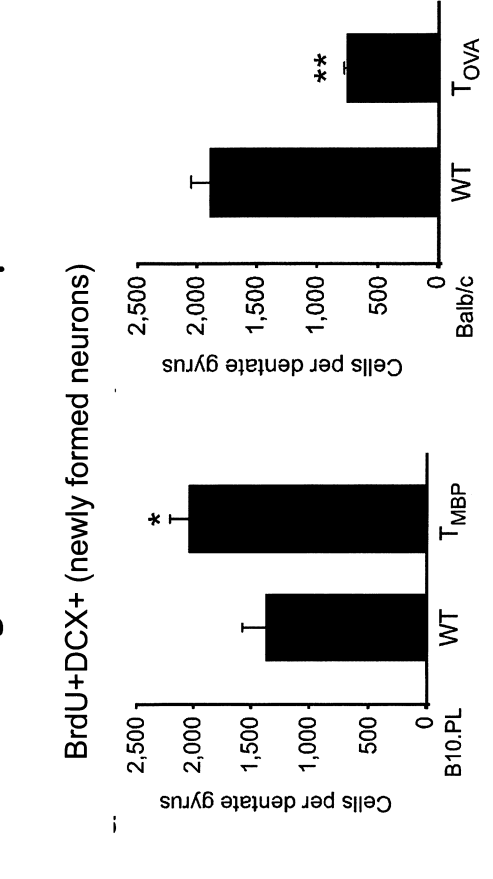
BrdU/DCX

5,000

4,000

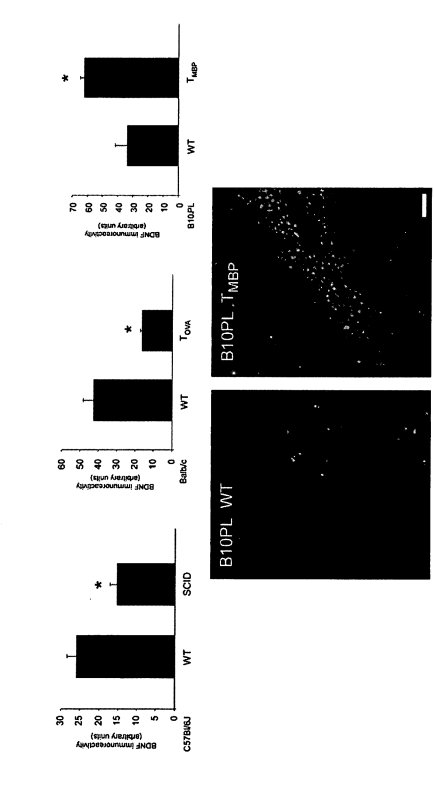
3,000

neurogenesis are CNS specific The T cells needed for adult



Ziv, Ron, Butovsky et al., Nature Neurosci. 2006

with hippocampal neurogenesis and learning abilities BDNF expression is T-cell related: Association



Ziv, Ron, Butovsky et al., Nature Neurosci. 2006



Leading Edge [New Foliology Select

of neurogenesis and indicate how these variables may be integrated during development and adulthood. This ation, migration, cell fate choice, and survival. Recent papers identify factors involved in each of these aspects recent work includes the report of a kinase that controls cell division in neural progenitors and a study suggesting that autoimmune T cells are positive regulators of neurogenesis. Other intriguing findings link planar cell polarity The formation of new neurons (neurogenesis) results from the complex interplay of many variables: cell proliferto neural tube development in zebrafish and the migration of neuroblasts in adult mice.

Autoimmunity Gives Neurogenesis a Lift

it suggests mechanisms by which age-related changes in the immune system could be linked to cognitive decline that recognize self-antigens have a bad reputation. However, new work by Schwartz, Kipnis, and colleagues argues Zivetal., 2006) for a more complex view of these much maligned cells. Their work suggests that, rather than always being detrimental, self-recognizing T cells can also support neurogenesis in adult mice if well-controlled. Previous in this case myelin basic protein) but is not restored by T cells that recognize a non-self-antigen. The presence of has been linked to neuronal activity in the hippocampus. One of the far-reaching implications of the study is that in humans. Future work may also establish the precise mechanisms by which T cells regulate microglia to foster an Given the link between autoimmunity and diseases such as multiple sclerosis, it is no wonder that T cells in the brain work from the Schwartz lab has shown that the recruitment of autoimmune T cells to sites of neuronal injury pronotes neural cell survival by altering the behavior of local microglia. This new report suggests that similar immune-based mechanisms may also operate during normal adult neurogenesis. In support of their argument, Ziv et al. (2006) demonstrate that neurogenesis is impaired in the hippocampus of immune-deficient mice. Moreover, unlike wild-type mice, neurogenesis in immune-deficient mice is not stimulated by enrichting the mouse's environautoimmune T cells also improves the performance of mice in the Morris water maze, a spatial memory task that ment. Remarkably, neurogenesis is restored by the introduction of autoimmune T cells that recognize a self-antigen environment that supports neuronal survival.

Ziv et al. (2006). Nature Neuroscience. Published online January 15, 2006. 10.1038/nn1629.

How can protective autoimmunity be boosted and developed as a therapeutic approach?

·Weak self-antigen (cryptic)

(Fisher et al., J. Neurosci., 2001;

·Altered self-antigen (APL)

(Hauben et al., J. Clin. Invest., 2001);

(Hauben, Gothilf et al., 2003) Dendritic cells

Self-reactive

T-cells

Boost

(Kipnis et al., PNAS 2000; Schori et al., PNAS 2001) Random copolymers

·Pharmacological blocking of Treg

regulate

Down

(Kipnis et al, 2004)

·Homeostasis-driven

(lymphopenia; Kipnis et al., E Proliferation of lymphocytes



Regulatory T-cells CD4+CD25+

Down

Poly (YE)

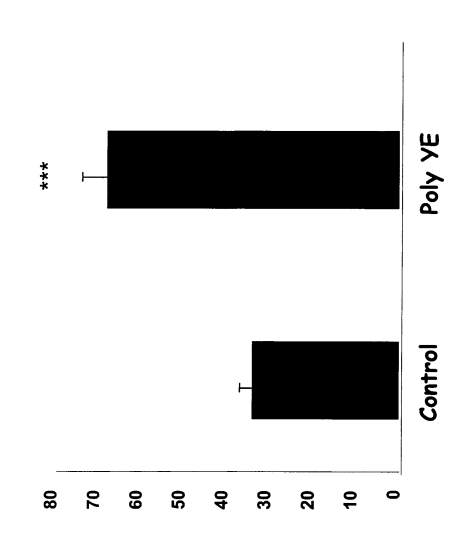


- 1. Random polymer
- Evokes strong immune response in mice
- Down regulates a subpopulation of regulatory T cells resulting in a speedy recruitment of the relevant autoimmune T cells

Proof of concept

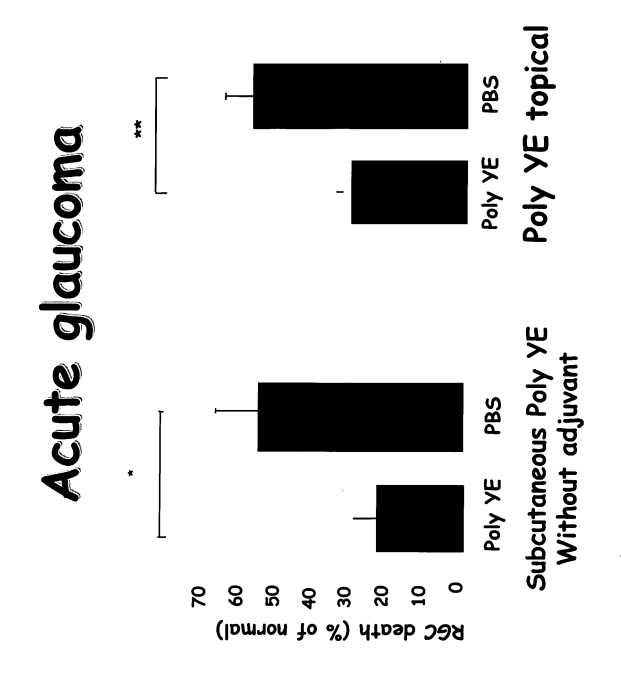
Sutamate toxicity

Proof of concept Slutamate toxicity

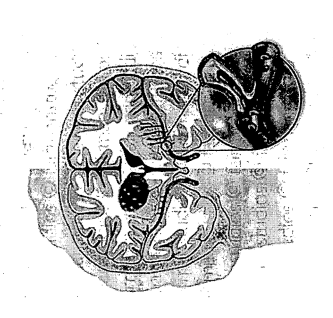


% RGC survival

Poly YE Neuroprotective effect in animal models of CNS disorders



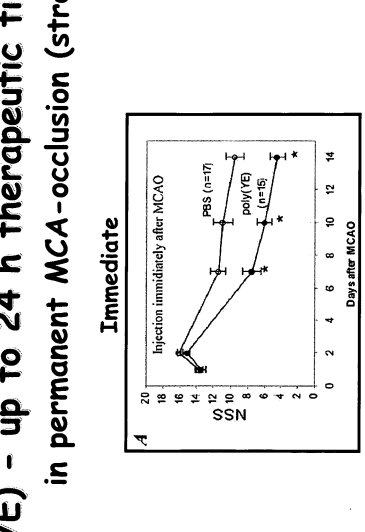
Stroke

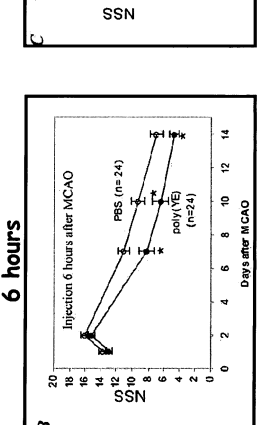


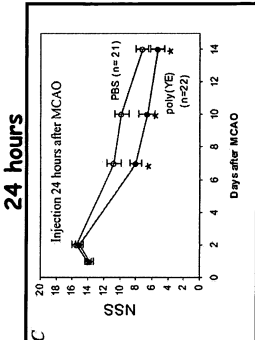
Middle cerebral artery occlusion (MCAO)

Poly(VE) - up to 24 h therapeutic time window

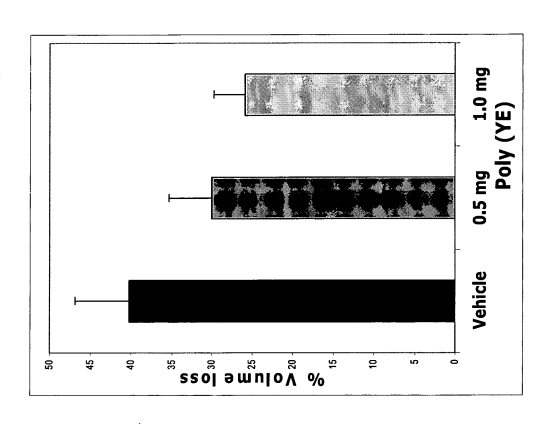
in permanent MCA-occlusion (stroke)



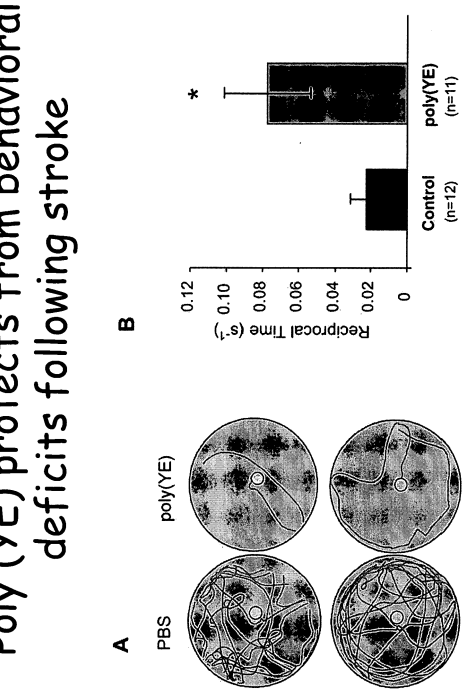




Poly(YE) decreases volume loss starting from the subacute phase

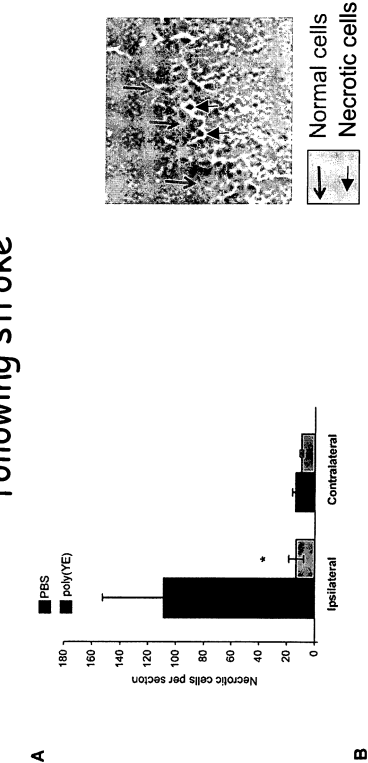


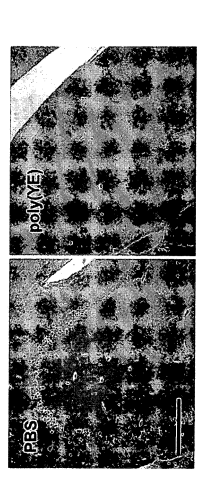
Poly (YE) protects from behavioral

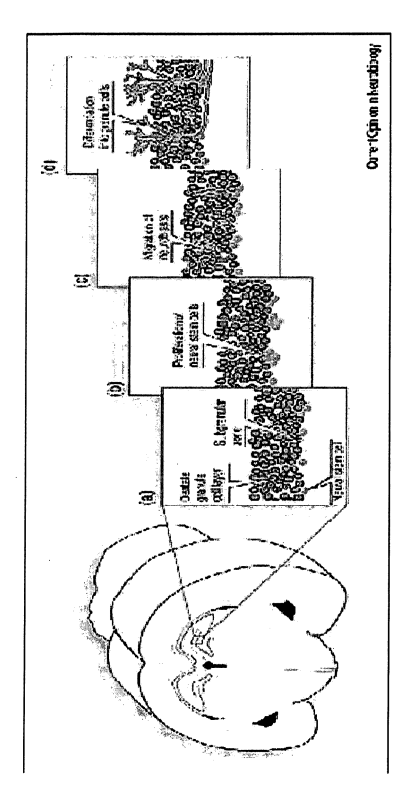


Poly (YE)- treated group learned and remembered the place of the platform unlike the control group

Poly (YE) attenuates hippocampal neuronal death following stroke

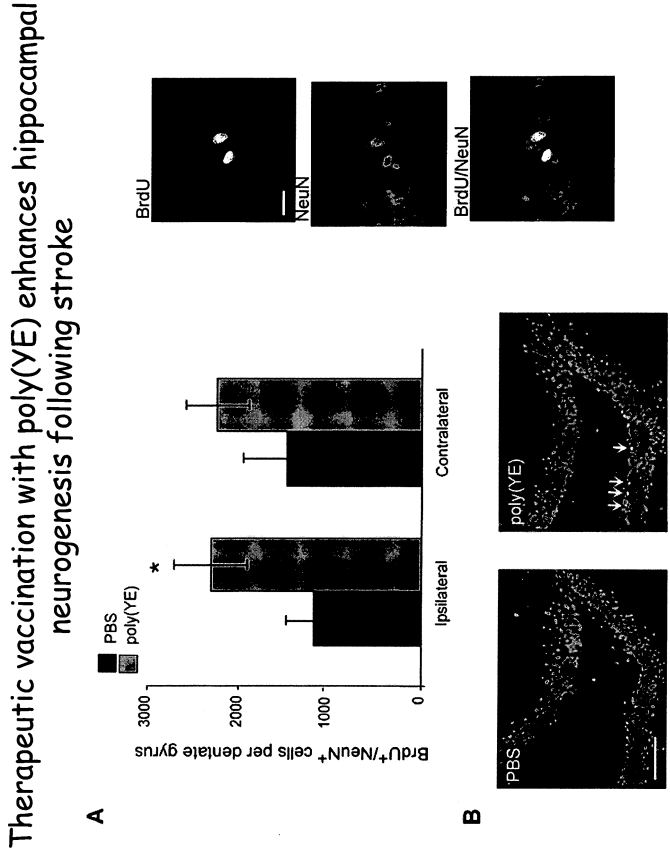


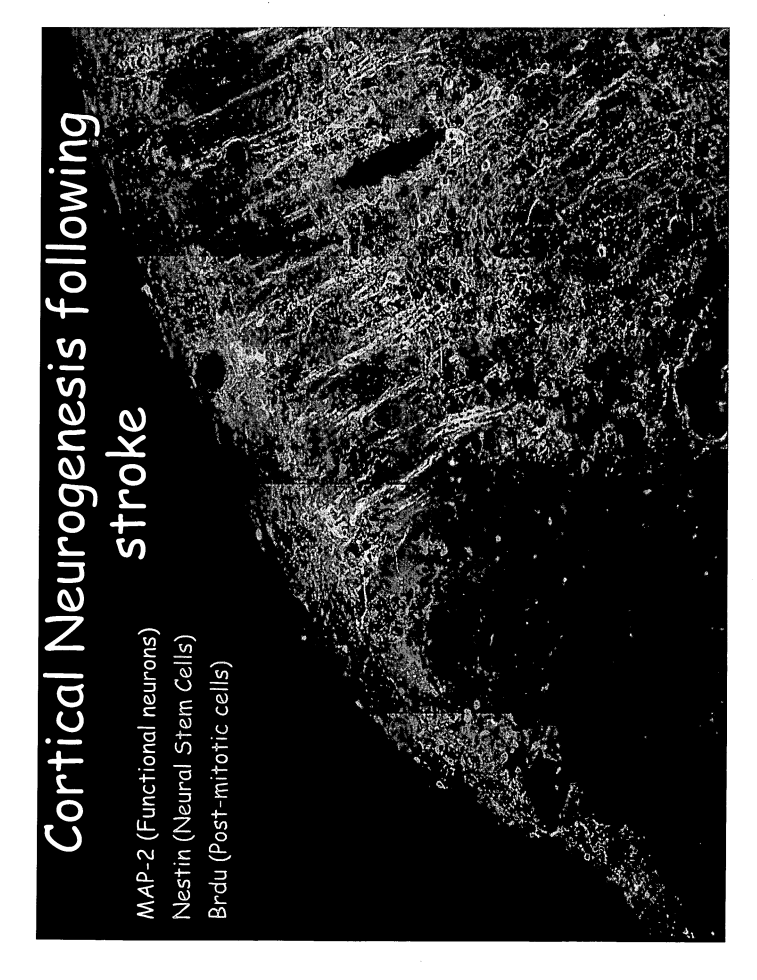




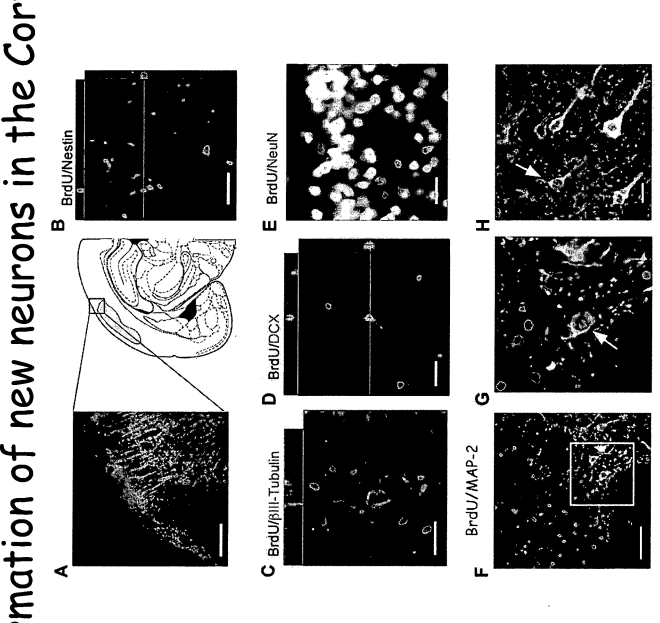
Hippocampal neurogenesis is increased after cerebral ischemia

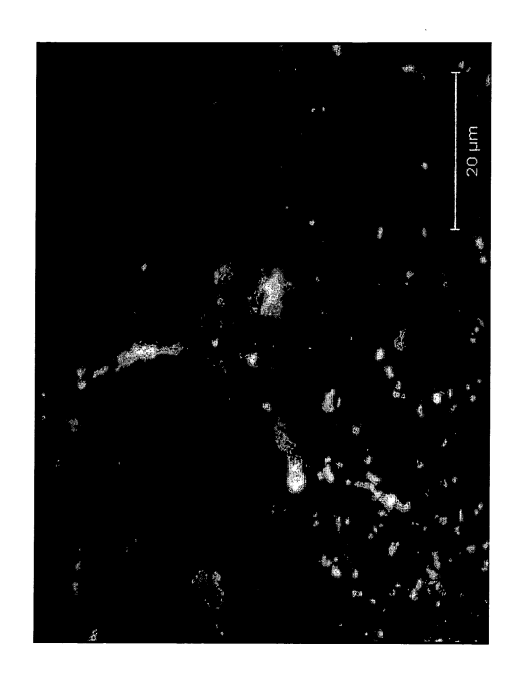
Nakatomi et al, Cell 2002 Kokaia & Lindvall, Cur Opin Neurobiol, 2003



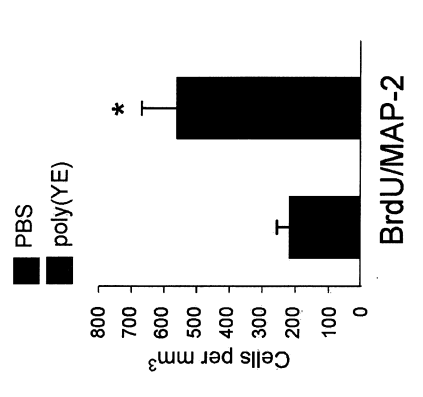


Formation of new neurons in the Cortex



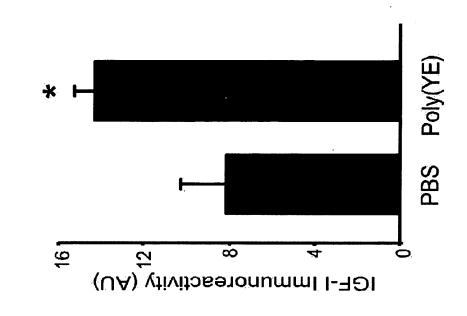


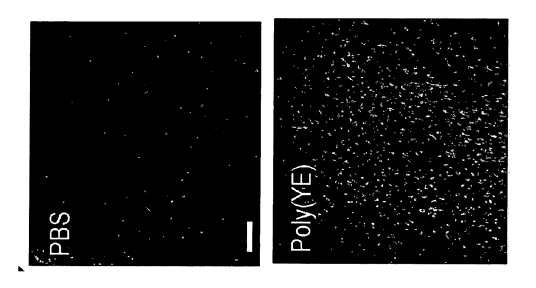
Poly(YE) augments cortical neurogenesis



Neurogenesis in a non-neurogenic area

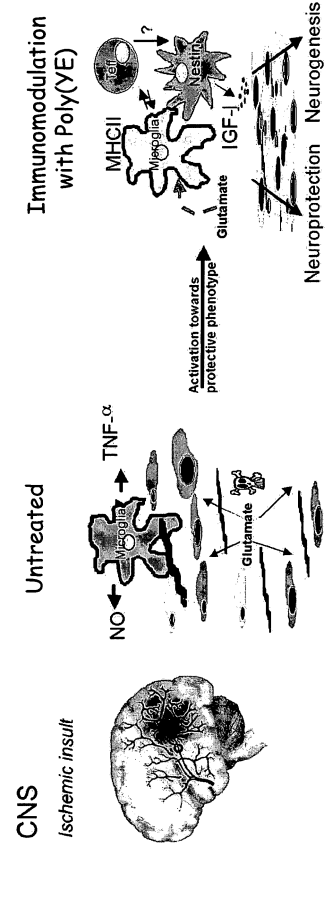
IGF-1 production is increased following Poly(YE)



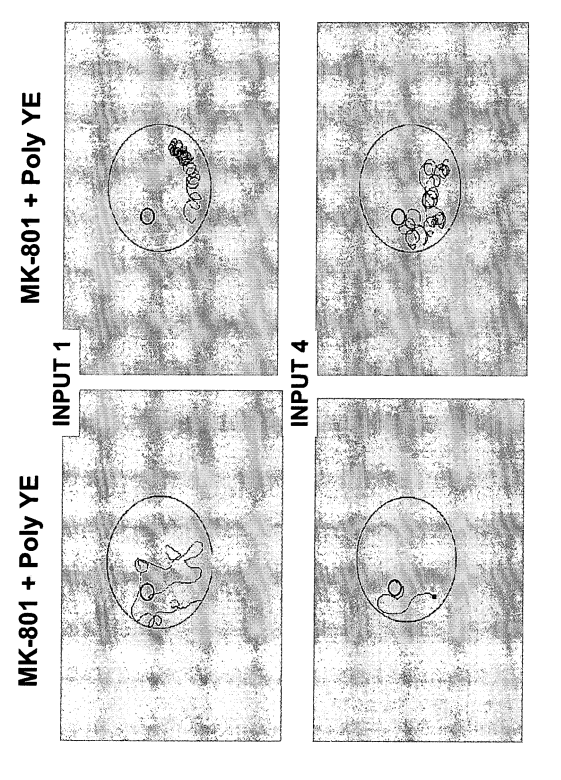


interactions in the ischemic brain after Putative model of the neuroimmune





Mental disorders



Protective immunity

Mechanism of action

Poly YE an immune modulator - augments the injury induced protective immunity by its effect on the local immune response

microglial activity to eliminate toxic elements from the injured environment and to secrete Poly YE restore tissue homeostasis - shapes trophic and growth factors, supportive of neuronal survival

Poly YE enhances tissue repair - creates environmental conditions supportive of neurogenesis